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- (54) METHOD FOR POLYMERIZING ETHYLENICALLY UNSATURATED MONOMERS BY DEGENERATIVE IODINE TRANSFER
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ABSTRACT

The invention pertains to a method for making a block or gradient final (co)polymer comprising a step of radically polymerizing a mixture of ethylenically unsaturated monomers in the presence of a) a radical precursor and b) an iodine atom-containing intermediate polymer or a mixture of iodine atom-containing intermediate polymers, wherein the icdine atom-containing intermediate polymer comprises at least 50 mole % of methacrylate monomers and is obtainable from a polymerization of ethylenically unsaturated monomers

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METHOD FOR POLYMERIZING ETHYLENICALLY UNSATURATED MONOMERS BY DEGENERATIVE IODINE TRANSFER

RELATED APPLICATIONS

[0001] This application claims priority based on European Patent Application No. 02078004.5, filed Jul. 23, 2003, and U.S. Provisional Patent Application No. 60/424,571, filed Nov. 7, 2002.

SUMMARY OF THE INVENTION

[0002] The invention portains to a method to propare a block or gradient co-polymer comprising a step of radically polymerizing a mixture of ethylemically unsaturated monomers.

BACKGROUND OF THE INVENTION

[0003] Controlled radical polymerization processes have received much attention over the past decade, because of the possibility to prepare new valuable polymeric materials based on standard conventional monomeric building blocks. such as block or gradient copolymers, and narrow-polydis-persity functional polymers, with narrower functionality distributions, that can be used in film-forming or crosslinking compositions, such as coating compositions, adhesives and printing ink formulations.

[0004] In the field of "living" radical polymerizations (polymerizations under radical conditions where termination processes of growing radicals are reversible, so that all initiated chains can, in principle, continue to grow as long as monomers and radicals are present), three technologies have been studied in depth:

[0005] nitroxide mediated polymerizations (NMP), which are based on homolytic scission of the covalent bond formed between a polymer radical and a nitroxide stable radical:

[0006] atom transfer radical polymerization (ATRP). in which a halogen (usually Cl or Br) atom is reversibly transferred between a Cu(I/II) complex and a polymer radical, in a process that ancompasses a redox cycle;

[0007] reversible addition fragmentation chain transfer (RAFT), which is a process wherein dithioester end groups can be rapidly and reversibly transferred between polymeric radicals.

[0008] Por all these systems, the living character was proven by a typically linear development of molecular weight with conversion, low polydispersities obtained in polymerizations, and the preparation of block copolymers by sequential addition of different monomers.

[0009] ATRP and RAFT especially have possibilities for numerous applications with respect to the types of monomers used. In ATRP, however, seidic conditions and seidic monomers create problems, while the necessary removal of the Cu-amine complexes from the polymers is an expansive and thus unwanted step. In RAFT, these limitations do not occur, but this method suffers in that the polymer chains formed contain dithioester and groups, which are strong chromophores. The chromophores can be destroyed by reaction with a nucleophile, but this nucleophile is not

always compatible with the functional groups that may be present in the chains. Such chromophore destruction comes at the expense of an extra reaction step and leads to low-molecular weight products in the polymer, which may be difficult to remove. Moreover, the dithiocster mediating compounds are expensive.

[0010] In general, RAFT based on dithio-ester compounds is assumed to be favourable in the rate of end group transfer (more transfer events, lower polydispersity, better control).

A lower polydispersibility is desirable when the polymer is used in e.g. film-forming or cross-linking compositions. A narrow molecular weight distribution in such applications leads to a good balance of pot life and curing speed, viscosity and network properties. Further, in the art it is assumed that the RAFT process is applicable to a broad range of monomers.

[0011] NMP has the disadvantage that expensive nitroxides are needed if the process is to be used at acceptably low temperatures for acrylates, whereas methacrylates have intrinsic problems due to the fact that disproportionation may occur as a side reaction.

[0012] Another type of living radical polymerization pro-Journal of Physical Organic Chemistry, Vol. 8, 306-315 (1995) and Macromoleoules, 28, 8051-8056 (1995)). In this process, as in the RAFT process, an iodine end group can transfer from one polymer to a radical end group of another polymer present in the polymerization system, the indine atom playing a similar role to that of the dithioester group in RAFT, the terminal lodine and the radical function exchanging between two polymer chains. A chain transfer agent with an indine atom is used as entry in the DIT process. Conventional judine chain transfer agents include alkyl or perfluoroalkyl iodides.

[0013] In Macromolecules, 28, 8051-8056 (1995), Macromolecules, 33(9), 3485 (2000), Macromolecules, 32(22), 7354 (1999), Macromolecules, 31(9), 2809 (1998) the DIT process was described for application with styrene monomers, for which it appears to work moderately well (polydispensities typically 1.5), and for application with halogenated vinyl monomers like vinyl chloride and fluorinated vinyl species.

[0014] DIT was attempted for acrylates, but the results showed less control (broader polydispersities>2, indicating a low rate of transfer between acrylate chains) (see: Macromolecules, 28, 8051-8056 (1995)). DIT processes with surplates have been disclosed in a few documents: however, these aim at obtaining iodine-functional chains instead of aiming at actual living polymerizations (see for instance U.S. Pat. No. 6,143,848). DIT processes to prepare block copolymers of acrylates and styrens have been reported in EP 947527, Macromolecules, 28, 2093 (1995), and Macromol. Rapid Commun., 2000, 21(13), 921.

[0015] A few attempts to obtain a degenerative iodine transfer process with methacrylates have been reported, but all of these suggest that methacrylate monomers cannot be used in an acceptable manner in conventional DIT processes using the conventional iodine-functional CTAs, which are applied in the DIT process for e.g. styrene (see for instance Macromolecules, 28, 8051 (1995)). High-molecular weight material is formed at the start of the polymerization, leading

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to high polydispersities, without the typical linear development of molecular weight with conversion, which is characteristic for a polymerization with a living character, being observed. Another comparison of DIT processes with styrene, acrylate, and methacrylate monomers was disclosed in B. Klumperman at the UNESCO School & South African IUPAC conference on Macromolecular and Materials Science, Mar. 29-31, 1999 and Apr. 10-12, 2000, which is to be found at http://www.sun.ac.za/unesco/PolymerED2000/Conf1999/Lectures1999/Klumpermac.pdf.

these respectively mere produced that DIT is not a suitable process for the polymerization of a mixture of monomers containing a large amount of methacrylate monomers. Theraffore, it has been assumed that DIT is not suitable for polymerizing methacrylates in a controlled manner, to chitain the hencefits that can be obtained in a "controlled (living) radical polymerization" process in the art. In U.S. Pat. No. 5,439,080 this is confirmed in Comparative example 2, when it was found that when MMA alone is used with a fluorinated alkyliaddide, only a homopolymer of MMA is produced and a block polymer with an iodide-functional perfluoropoly-other is not produced.

[0016] Since methacrylatos are a very important class of monomers for many applications, with high- and low-polarity candidates and various functional side groups available, with high-Tg backbones with good chemical durability being formed, the inability to use methacrylates is a scrious problem in DIT technology, and a need exists for an effective DIT method allowing the production of polymers based on methacrylates with structures that benefit from the advantages of the living nature of the polymerization process.

[0017] It is an object of the present invention to obtain a DIT process that can be used for (co)polymentzing methorylate monomers as in a living radical polymerization process. According to the present invention, it was found that the DIT process can be performed in a very favourable way using methorylates, if the proper start-up process is chosen. We have found that the DIT process can be adapted for the polymerization of compositions of predominantly methorylate monomers, to find a high transfer rate of foidize atoms between methorylate chain ends, leading to better control than a DIT process using styrend or acrylate monomers, to low polydisponsities closely resembling those of ATRP or RAFT processes, molecular weights increasing with conversion, and the apportunity to prepare well defined block and gradient copolymers.

DETAILED DESCRIPTION OF THE

[0018] We have found a novel method wherein iodinefunctional methacrylate polymers are obtained, leading to a DIT process proceeding with a high level of control.

[D019] More specifically, the invention pertains to a method for making a block or gradient final co-polymer comprising a step of radically polymerizing a mixture of ethylenically unsaturated monomers in the presence of a) a radical precursor and b) an iodine atom-containing intermediate polymer or a mixture of iodine atom-containing intermediate polymer-comprises at least 50 mole % of methacrylate monomers and is obtainable from a polymerization of ethylenically unsaturated monomers.

[0020] For clarity purposes the terms "intermediate polymer" and "final polymer" are introduced in this description. However, it has to be understood that the final polymer can be reacted further and that the polymerization steps leading to the intermediate and the final polymer need not follow each other directly, but that there may be (a) further processing or reaction step(s) between the intermediate and fleat polymerization).

[0021] The iodine atom-containing intermediate polymers to be formed initially (forming the "first block" if a block polymer is to be made) should predominantly comprises monomers of the methacrylate type, i.e. the polymer comprises at least 50 mole % of methacrylate monomers (in this application methacrylate monomers include methacrylic acid or exters, amides or anhydrides thereof, or methacryle lonitrile). This results in the end group of the intermediate polymer being predominantly of the iodine-containing methacrylate type.

[0022] Such indine-containing methacrylate and group has

[0023] wherein P stands for polymer and CX is an acid, anhydride, ester, amide, or nitrile group.

[0024] More preferred is a composition wherein the intermediate polymer comprises more than 70 mole % of methacylate monomers; most preferred is a composition wherein the intermediate polymer comprises more than 90 mole % of this intermediate polymer is less than 20,000 and more than this intermediate polymer is less than 20,000 and more than 5,000 and more than 1,000, most preferably less than 10,000, most preferably less than 5,000 and more than 1,000. In this respect the use of the term intermediate oligomer instead of intermediate polymer should be taken into consideration as well. The todine atom-centaining intermediate polymer may additionally comprise other (colmonomers which may be of the (meth)serylate, styrens, vinyl ester, and malesto type.

[0025] The polymerization step leading to the final polymer can be performed with the fadine atom-containing intermediate polymer and a monomer or a mixture of monomers, the composition of which may be different from the composition of the indine atom-containing intermediate polymer, in the presence of a radical procursor.

[0026] Examples of suitable mixtures for making the final and intermediate polymers comprise monomers selected from, e.g. scrylic acid, methacrylic acid, methyl scrylate, chyl scrylate, butyl acrylate, methyl methacrylate, ethyl methacrylate, butyl methacrylate, glycidyl acrylate, butyl methacrylate, glycidyl acrylate, 2-hydroxychyl acrylate, 2-ethylhoxyl acrylate, potassium methacrylate, atearyl methacrylate, 2-dimethylamino)chyl methacrylate, atearyl methacrylate, 2-hydroxychyl methacrylate, butyl methacrylate, lauryl methacrylate, diacetone acrylamide, acrylamide, acrolein, methacrylatinde, methacrylate, styrene, siphamethyl syrene, vinyl nchane; scoproponyl-syrene, siphamethyl syrene, vinyl nchane; scoproponyl-

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isopropylidene isocyanatobenzene (mTMI from Cytec), chlorostyrene, butadiene, isoprene, chloroprene, maleic acid, maleimide, methyl maleate, ethyl maleate, propyl malcate, butyl malcate, calcium maleate, allyl malcate, 2-ethylhexyl maleate, octyl maleate, maleio hydrazide maleic anhydride, fumaric acid, methyl fumarate, ethyl fumarate, propyl fumarate, butyl fumarate, sodium fumarate, fumaronitrile, fumaryl chloride, vinyl acctate, tertdecanoic acid ethenyl ester (VcoVa® 10, ex Resolution Performance Products), neononanoic acid ethenyl ester (VcoVa® 9, cx Resolution Performance Products), vinyl pivalate, and the like. It is not necessary that methacrylate monomers are predominantly present in the mixture for the final polymer, and this mixture may even be free of methacrylic monomers. If the monomers selected for this final polymerization step are predominantly of the methacrylate type, the process can then be repeated to form a third or further block, or gradient copolymers can be obtained, as is obvious to the skilled person.

[0027] The weight ratio of the indine atom-containing intermediate polymer to the monomers being polymerized to form the final polymer may vary from 1:100 to 100:1.

[0028] Optionally, the amounts are selected so that the mole ratio incline atom-containing informediate polymer b): radical precursor a) is 50.1 n, more preferably >0.5 n, wherein n stands for the number of radicals effectively generated per molecule of radical precursor (i.e. taking into account the initiator efficiency).

[0029] The final polymerization process can take place in batch mode or semi-batch mode, with feeding of several of the reactants to the vessel, or in a process comprising a continuous step. The iodine-functional methacrylate intermediate polymers of the DIT type can be transferred and used as a CTA in the final polymerization process according to the invention.

[0030] The final polymerization can be carried out in bulk, in an organic solvent or in an aqueous medium, in a suspension polymerization process, an emulsion polymerization process, a dispension polymerization process, or a mini-emulsion polymerization process.

[0031] The iodine atom-containing intermediate polymer is made from the monomer mixtures as mentioned above for the final polymer, provided that the amount of methacrylic monomers is so selected that at least 50 mole % of the menomers are of the methacrylate type. The polymerization leading to the intermediate polymer is performed in the presence of \(^1_2\) or an effective iodine chain transfer agent (CTA) d) and a radical precursor c). The term iodine as used in this application includes \(^1_2\) and \(^1_2\)-precursors (i.e. compounds that can form \(^1_2\) in situ), such as N-iodosuccinimide, ICI, IBr, and \(^1_2\)-indime chain transfer agents and iodine are used in combination with a small amount of radical precursor to obtain a methacrylate intermediate polymerization

[0032] When molecular iodine is used, the lodine radical released after one iodine is abstracted by a polymer radical does not reinitiate a methacrylate polymerization, but recombines with another polymer radical. The net effect of molecular iodine is the functionalization of two polymer chains with an iodine end group, in an identical way U. Polym. Sct. Polym. Lett. Ed., Vol. 14, 499-502 (1976)). In

terms of the intermediate polymerization process, it implies that slightly more than 1 equivalent of imitiating radicals should be generated, compared to the iodine atoms present in the systems as molecular iodine. The mole ratio 1₂: radical precursor of preferably is between 0.05 n and 0.5 n, more preferably between 0.1 n and 0.5 n, and most preferably between 0.25 n and 0.5 n, wherein n has the previously given meaning. This route to obtaining the low-molecular weight intermediate 1-terminated polymers is very effective, and is a preferred subordinent.

[0033] An example of an effective todine CTA (rapidly allowing transfer of the iodine to methacrylate polymer radicals) is a sulfneyl todide, e.g. p-toluens sulfonyl iodide. Whon such an iodine CTA is used, preferably the mole ratio of iodine CTA: radical precursor d) is >0.1 n, more preferably >0.5 n, wherein n stands for the number of radicals affectively generated per molecule of radical precursor.

[0034] The most preferred process according to the present invention is the one wherein iodine is used.

[0035] It was found to be advantageous sometimes to perform the polymerization stop(s) of the method according to this invention in the presence of an epoxide-containing compound, especially the DIT polymerization to form the intermediate polymer. In the presence of epoxide-containing compounds the polydispensity obtained is narrower, leading to coating compositions with a good balance of pol life and curing speed, viscosity and network properties. Similarly, such epoxide-containing compound can also be used in the synthesis of the final polymer. Such compounds are preferably selected from glycidyl ethers, glycidyl esters, and cycl-aliphatic epoxides. Suitably glycidyl-containing monomers are used in the preparation of the lodine atom-containing polymer, such as glycidyl methacrylate, glycidyl aerylate and the glycidyl ester of neodecancie acid (Cardura® E10). The method according to the invention is preferably performed such that the mole ratio epoxide iodice atom-containing polymer b) is >0.01, preferably >0.05.

[0036] Although the essence of the present invention is the radical polymerization of a mixture of citylenically unsaturated monomers in the presence of a) a radical procursor and b) an iodine atom-containing polymer or a mixture of iodine atom-containing polymers, the invention also portains to the two-step procedure for making a block or gradient co-polymer comprising a step of radically polymerizing a mixture of citylenically unsaturated monomers to give iodina atom-containing polymers h) comprising at least 50 mole % of methacrylate monomers in the presence of c) a radical precursor and d) l_a or a iodine chain transfer agent, followed by a step of radically polymerizing a mixture of ethylenically unsaturated monomers in the presence of a) a radical procursor and b) the iodine atom-containing polymer of the first step. The second step may be started before the first step is fully ended.

[0037] Also in this two-step procedure the mole ratio sulfonyl iodide: radical procursor is > 0.1 a, wherein a stands for the number of radicals effectively generated per molecule of radical precursor, or wherein the mole ratio I₃: radical procursor is between 0.05 a and 0.5 a.

[0038] The polymerization process(cs), i.e. the process leading to the intermediate polymer and/or the process leading to the final polymer, of the monomers take(s) places

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at temperatures of less than 130° C., preferably less than 110° C., even more preferably less than 90° C., most preferably less than 90° C., most mild initiating radicals is used as a radical precursor. Mild initiating radicals are for example C-centered radicals, preferably C-centered radicals obtained from the decomposition of ago radical initiators. Initiators of the szo-type include architecturymonitric (AIBN) or azobisystemotivements.

[0039] As indicated above, the final polymer may be reacted further. It may be advantageous to modify the indine and groups-containing polymers formed in the process according to the invention in a process after or during the polymerization. The end groups can be modified, e.g., by elimination, cyclization, atom transfar, redox, or substitution reaction, optionally under removal of todine and/or low-molecular weight todine-containing compounds. More specifically, the removal of the iodine atom may be done by any mechanism known to the skilled person such as a nucleophilic reaction, by heating of the polymer (leading to homolytic scission but also to elimination products), or by coation with a radical generating compound. It may be favourable to perform such thermal or radical creations in the presence of reducing conditions to scavenge loddine radicals or I₂ formed.

[0040] The nucleophilic reaction can suitably be performed using mercaptopropionic acid in combination with a base. The radical-generating compound may be a percaide, for example H₂O₂. A radical reaction under reducing conditions can be performed by heating the iodips-functional polymers in the presence of sodium formaldehyde sulfoxylate.

[0041] The final polymer prepared by the method according to this invention can be used in film-forming or cross-linking compositions such as coating compositions, adhasivea, and (printing) this formulations. Block or gradient copolymer products made by the method according to the invention can be used as rheology additives, surfactants, dispersants, adhesion promotors and/or flow improvement additives.

[0642] Prefered costing compositions include costing compositions that can be used in automotive and general industrial coating applications.

[0043] The invention is further illustrated with the following examples.

[0044] All reported molecular weights were determined by size exclusion chromatugraphy (RI detection) and are expressed in polystyrene equivalent molecular weights, unless specified otherwise. Low-molecular weight materials (M<500), originating e.g. also from initiator in cage recombination products and optionally added low-molecular weight spoxides, were not taken into account for this determination of the main polymer cavelope in the molecular mass distribution. The following abbreviations are used: MMA-methyl methacrylate, GMA-glycidyl methacrylate; BA-butyl acrylate, BMA-butyl methacrylate, HEMA-by-droxychyl methacrylate, SEC=size exclusion chromatography. Mn-number average molecular weight, Mv-Z-average molecular weight. The polydispersity of the polymers was determined as the ratio MyMn.

[0045] The effect of copolymerizing methacrylates and acrylates on molar mass distribution, wherein the molar

mass and the polydispersity are taken as a measure of the extent of control, is illustrated by the following examples. It can be seen that coatrol is lost if the fraction of acrylate monomers becomes too high. These experiments show that upon using MMA/BA ratios with a good control gradient, copolymers are obtained due to the higher polymerization reactivity of MMA monomer vs. BA monomer, as is known in the art.

[0046] In general, all copolymenizations carried out under conditions which allow for composition drift will produce gradient copolymers under the conditions of the DIT process in this invention.

EXAMPLE 1

[0047] A reaction vessel was charged with 21.25 g of MMA, 5.32 g of BA, 20 g of buyl acctate, 1.54 g of I₃, and 2.25 g of 2,2-ambis(2,4-dimethylvalerositrile), degassed, and maintained under a nitrogen atmosphere. The temperature was raised to 67° C, and the reaction was continued for 8 h. After 1 h and 40 min, the brown iodine colour was observed to fade, an observation that coincided with a steep increase of conversion with time, after the inhibition period in the presence of molecular iodine. SEC analysis of a sample taken after this period (conversion 88%) yields an Mn of 2,325 and an Mw/Mn of 1.31, Mz/Mw 1.25.

EXAMPLE 2

[0048] A reaction vessel was charged with 15.96 µg of MMA, 10.66 g of BA, 20 g of butyl acetale, 1.33 g of L₂, and 2.26 g of 2.2°-axobis(2,4-dimethylvaleronitrile), degassed, and maintained under a nitrogen atmosphere. The temperature was raised to 64° C., and the reaction was continued for 8 h. After 2 h the brown iodine colour was observed to fade, an observation which coincided with a steep increase of conversion with time, after the inhibitive period in the presence of molecular iodine. SEC analysis of a sample taken after this period (conversion 82%) yields an Mn of 2,210 and an Mw/Mn of 1.48, Mz/Mw 1.47.

EXAMPLE 3

[0049] A reaction vessel was charged with 10.62 g of MMA, 15.96 g of BA, 20 g of buryl acetate, 1.53 g of 7₄, and 2.207 g of 7_{2.2}-azobis (2.4-dimethylvalcronifitle), degassed, and maintained under a nitrogen atmosphere. The temperature was raised to 64° C, and the reaction was continued for 20 h. After 2 h, the brown inotine colour was observed to fade. SEC analysis of a sample taken after this period (conversion 95%) yields an Mn of 2,430, an Mw/Mn of 2.04, and an Mz/Mw of 2.16.

COMPARISON EXAMPLE 4

[0050] A reaction vessel was charged with 5.31 g of MMA, 21.26 g of BA, 20 g of butyl accrate. 1.52 g of 1₂, and 2.267 g of 2,2°-azobia(2,4-dimethylvalaronitrile), degassed, and maintained under a nitrogen atmosphere. The temperature was raised to 64° C., and the reaction was continued for 20 h. SEC analysis of a sample taken after this period (convestion 95%) yields an Mn of 2,580, an Mw/Mn of 2.17, and an Mz/Mw of 2.28.

COMPARISON EXAMPLE 5

[0051] A reaction vessel was charged with 25 g of BA, 20 g of butyl acetate, 1.67 g of I₂, and 2.267 g of 2,2'-azobis(2,

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4-dimethylvaleronitrile), degassed, and maintained under a nitrogen atmosphere. The temperature was raised to 64° C., and the reaction was continued for 4 h. SEC analysis of a sample taken after this period (conversion 93%) yields an Mn of 2,300, an Mw/Mn of 1.92, and an Mz/Mw of 2.07.

EXAMPLE 6

[0052] A reaction vessel was charged with 11.01 g of MMA, 11.00 g of BA, 8.06 g of HEMA, 20 g of butyl acetate, 1.523 g of la, and 2.26 g of 2.2'-azobis(2,4-dimethylvaleronitrile), degassed, and maintained under a nitrogen atmosphere. The temperature was raised to 64° C., and the reaction was continued for 7 h. SEC analysis of a sample taken after this period (conversion 98%) yields an Mn of 4,800, an Mw/Mn of 1.26, and an Mz/Mw of 1.31. This example shows that monomers with functional side groups may also be used.

EXAMPLE 7

[0053] To a reaction vessel maintained at 55° C. a solution consisting of 36.49 g of BMA, 6.57 g of GMA, 3.69 g of L, and 5.42 g of 2,2'-azobis(2,4-dimethylvaleronitrile) and 16 g of Dowanoid PM (glycol other solvent ex Dow Chemical Co.) was fed over a period of 2 h, to obtain a first BMAco-GMA block of Mn 2,880, Mw/Mn 1.25, Mz/Mw 1.19.

EXAMPLE 8

[0054] A reaction vessel was charged with 11.02 g of MMA, 11.01 g of BA, 20.06 g of buyl acetate, 1.52 g of I₂, and 2.26 g of 2.2° azobis(2.4-dimethylvaleronitrile), degassed, and maintained under a nitrogen atmosphere. The temperature was raised to 64° C., and the reaction was monitored until the dark indine colour started to fade (conversion 9%). At this point, 5.29 g of methacrylic acid and 2.75 g of Cardura B-10 were added in one shot and the reaction was continued for 4 h. SEC analysis of a sample taken after this period (conversion 99%) yields an Mn of 2,928, an Mw/Mn of 1.32, and an Mz/Mw of 1.28.

EXAMPLE 9

[0055] To the polymer of Example 7, a mixture of 27.9 g of HEMA and 0.469 g of 2,2'-azobis(2,4-dimethylvalaronitrile) was added, together with 10 ml of Dowanol@ PM, at a temperature of 64° C. After 4 h, a block copolymer was obtained with Mu 2,430, Mw/Mn 1.46, Mz/Mw 1.31.

EXAMPLE 10

[0056] A reaction vessel was charged with 30 g of MMA. 20 g of butyl acetate, 1.516 g of lo, and 2.259 g of 2,2'-azobis(2,4-dimethylvalcronitrile), degassed, and maintained under a nitrogen atmosphere. The temperature was raised to 66° C., and the reaction was continued for 4.5 h. At this point, the conversion was 81%, the Mw was 3,000 (Mz/Mw 1.26), and a mixture of 5.01 g of HEMA, 5.08 g of BMA, 6 g of butyl acctate, and 0.0097 g of 2,2'-azobis(2, 4-dimethylvaleronitrile) was added, and the reaction was continued for another 3.5 h. The final block copolymer had an Mw of 3.580, Mz/Mw 1.33).

[0057] A reaction vessel was charged with 30.05 g of MMA, 20.05 g of butyl acetate, 1.52 g of I2, 2.26 g of

2,2'-azobis(2,4-dimethylvalcronitrile), degassed, and maintained under a nitrogen atmosphere. The temperature was raised to 65° C., and the reaction was continued for 6.5 h. SEC analysis of a sample taken after this period (conversion 97%) yields an Mn of 2,881, an Mw/Mn of 1.14, Mz/Mw

[0058] Three days later, a mixture of 60 g of BMA, 35.0 g of buyl acctate, and 0.80 g of 2,2'-azobis(2,4-dimeth-ylvaleronitrile) was slowly added to this block over a period of 3.5 h at 65° C., and polymerized for an additional 2 h to obtain a block copolymer with Mn 7,200, Mw/Mn 1.67, Mz/Mw 1.47.

EXAMPLE 12

[0059] A reaction vessel was charged with 30.0 g of MMA, 20 g of butyl acctate, 2.54 g of I₂, 3.83 g of 2,2-azobis(2.4-dimethylvaleronitrile), and 2.33 g of Cardura@ E-10 (versatic acid glycidyl ester, ex Resolution), degasted, and maintained under a nitrogen atmosphere. The temperature was raised to 64° C., and the reaction was continued for 5 h. SEC analysis of a sample taken after this period (conversion 95%) yields an Mn of 2,990, an Mw/Mo of 1.20, Mz/Mw 1.16.

EXAMPLE 13

[0060] To the polymer of Example 12 was added a mixture of 2.57 g of BMA, 2.5 g of butyl acetate, and 0.257 g of 2,2'-azobis(2,4-dimethylvaleronitrile). The reaction was continued for an additional 2 h (conversion 99%), to yield a block copolymer with Mn 3,263, Mw/Mn 1,21, Mz/Mw

EXAMPLE 14

[0061] A reaction vessel was charged with 30.05 g of MMA, 20.03 g of butyl acetate, 2.54 g of 12, 3.79 g of 2,2'-azobis(2,4-dimethylvaleronitrile), degased, and maintained under a mitrogen atmosphere. The temperature was raised to 64° C., and the reaction was continued for 7 h. SEC analysis of a sample taken after this period (conversion 97%) yields an Mn of 1710, an Mw/Mn of 1.16, Mz/Mw

[0062] To this block was added a mixture of 15.04 g of BMA, 10.06 g of butyl acctate, and 0.27 g of 2,2-arobis(2, 4-dimethylvaleronitrile) at a feeding rate of 0.187 ml/min. The polymerization of the second block was continued for a total time of 4.5 h to yield a block copulymer with Mn 2,390, Mw/Mn 1.31, Mz/Mw 1.38.

[0063] Next to the I2 route, a good start to a DIT process with methacrylates can also be made if an iodine CTA, such as tosyl iodide, with a high chain transfer coefficient for methacrylates (CTC>1) is used.

[0064] The foregoing description of the present invention is merely illustrative thereof, and it is understood that variations and modification can be made without departing from the spirit or scope of the invention.

1. A method for making a block or gradient final (co)polymer comprising a first step of radically polymerizing a mixture of chylenically unsaturated monomers to an iodine atom-containing intermediate polymer, wherein the todine atom-containing intermediate polymer comprises at least 50

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mole % of methacrylate monomers, in the presence of a radical procursor and an In or an iodine chain transfer agent, followed by a second step of radically polymerizing a mixture of ethylenically unsaturated monomers in the presonce of a radical precursor and the iodine atom-containing intermediate polymer of the first step.

- 2. The method according to claim 1 wherein the mole ratio of the iodine atom-containing intermediate polymers to the radical procursor of the second step is greater than 0.1 n, wherein n stands for the number of radicals effectively generated per molecule of radical precursor.
- 3. The method according to claim 1 wherein the polymerization occurs at a temperature lower than about 130° C.
- 4. The method according to claim 3 wherein the temperature is lower than 110° C.
- 5. The method according to claim 3 wherein the temperature is lower than 90° C.
- 6. The method according to claim 3 wherein the temperature is lower than 70° C.
- 7. The method according to claim 1 wherein the polymerization in the first and second steps are performed in the presence of an epoxide-containing compound.
- 8. The method according to claim 7 wherein the mole ratio of the apoxide to the fodine atom-containing intermediate polymer is greater than 0.01.
- The method according to claim 8 wherein the mole ratio of the epoxide to the judine atom-containing intermediate polymer is greater than 0.05.
- 10. A method for making a block or gradient final (co)polymer comprising a step of radically polymerizing a mixture of ethylenically unsaturated monomers in the presence of a radical precursor and an iodine atom-containing intermediate polymer or a mixture of iodine atom-containing intermediate polymers, wherein the iodine atom-containing intermediate polymer comprises at least 50 mole % of methacrylate monomers and is obtainable from a polymerization of ethylenically unsaturated monomers.
- 11. The method according to claim 10 wherein the mole ratio of the iodine atom-containing intermediate polymer to the radical precursor is greater than 0.1 n, wherein n stands for the number of radicals effectively generated per molccule of radical procursor.
- 12. The method according to claim 10 wherein the temerature during the polymerization step is lower than about 130° C.
- 13. The method according to claim 12 wherein the temperature is lower than 110° C.
- 14. The method according to claim 12 wherein the temperature is lower than 90° C.
- 15. The method according to claim 12 wherein the temperature is lower than 70° C.
- 16. The method secording to claim 10 wherein the polymerization step is performed in the presence of an epoxidecontaining compound.
- 17. The method according to claim 16 wherein the mole ratio of the epoxide to the todine atom-containing intermediate polymer is greater than 0.01.

- 18. The method according to claim 16 wherein the mole ratio of the epoxide to the iodine atom-containing intermediato polymer is greater than 0.05.
- 19. A method according to claim 10 wherein the iodine atom-containing intermediate polymer is obtainable by polymerization of a mixture of ethylenically unsaturated monomers comprising at least 50 mole % of methacrylate monomers in the presence of a radical precursor and an iodine or an iodine chain transfer agent.
- 20. The method according to claim I wherein the mole ratio of the I2 to the radical precursor of the first step is between 0.05 n and 0.5 n, wherein n stands for the number of radicals effectively generated per molecule of radical
- 21. The method according to claim 1 wherein the iodine chain transfer agent is sulfonyl iodidc.
- 22. The method according to claim 21 wherein the mole ratio of the sulfonyl lodide to the radical precursor of the first step is greater than 0.1 o, wherein a stands for the number of radicals effectively generated per molecule of radical
- 23. A method according to claim 1 wherein the iodine atom-containing intermediate polymer has a molecular weight of less than 10,000.
- 24. A method according to claim 1 further comprising a third step of removing the iodine atom in the final polymer.
- 25. The method scoonling to claim 24 wherein the iodine atom is removed by nucleophilic reaction, by heating, or by reaction with a radical-generating compound, optionally under reducing conditions.
- 26. A block or gradient (co)polymer obtainable by the method of claim 1.
- 27. A film forming composition comprising the block of gradient (co)polymer of claim 26.
- 28. A coating composition, adhesive or ink formulation comprising the block or gradient (co)polymer of claim 26.
- 29. An automotive or industrial coating composition comprising the the block or gradient (co)polymer of claim
- 30. A rheology additive, surfactant, dispersant, adhesion romoter or flow improvement additive comprising the block or gradient final (co)polymer of claim 26.
- 31. A block or gradient (co)polymer obtainable by the method of claim 10
- 32. A film forming composition comprising the block or gradient (co)pulymer of claim 31.
- 33. A coating composition, adhesive or ink formulation comprising the block or gradient (co)polymer of claim 31.
- 34. An automotive or industrial coating composition comprising the the block or gradient (co)polymer of claim
- 35. A theology additive, surfactant, dispersant, adhesion promoter or flow improvement additive comprising the block or gradient final (co)polymer of claim 31.